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Comparative dissolution of electrospun Al₂O₃ nanofibres in artificial human lung fluids

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Abstract

Sub-micron sized alumina fibres were fabricated by electrospinning and calcination of a polymer template fibre. In the calcination step, different controlled temperature heating cycles were conducted to obtain fibres of different crystalline structures. Their biodurabilities were tested at pH 7.4 with lung airway epithelial lining fluid or serum ultrafiltrate (SUF) and at pH 4.5 with macrophage phagolysosomal simulant fluid (PSF). Potential to generate free radicals was tested in vitro. Through the variation in the soak temperature from 650 °C to 950 °C (experiments S650-S950), the heating protocol affected the morphological characteristics, crystal structure, surface area, and density of the alumina fibres while their dissolution half-times were not significantly affected in SUF or PSF. Fibre samples formed at different heating ramp rates (experiments R93– R600) showed significant variation in the dissolution rates with the highest ramp rate corresponding to the highest dissolution rate. Thus, by increasing the calcination temperature ramp rate the alumina fibres may be produced that have reduced biodurability and lower inflammogenic potential. The fibres with the highest dissolution rated had the least aluminium content. The solubility half-times of the alumina fibres were shortest for fibres calcined at the fastest temperature ramp rate (though soak temperature did not have an effect). The ramp rates also affected the aluminium content of the fibres suggesting that the content may affect the structural strength of the fibres and control the dissolution.

Introduction

Sub-micron and nanoscale (1–100 nm) diameter alumina fibres are of interest for a number of potential applications such as catalyst support structures, reaction substrates, filtration, and sensors. ^{1–5} The small diameters of the fibres, and the small dimensions of nanomaterials in general, enhance material properties through high surface areas per unit mass and enhance transport properties through slip and molecular flow effects. ⁶ The enhanced properties make

these materials attractive to industries for development of new advanced fibre materials. Sub-micron and nanoscale diameter fibres are working their way into the fabrication processes of common materials such as filter media for air filtration.⁷

Environmental, health, and safety concerns of nanotechnology have received attention from organizations such as the National Academies. The manufacture, use, and disposal of engineered nanomaterials may lead to exposure of humans to the nanomaterials. Because these materials are relatively new and their biological behaviour may differ from that of the bulk material with the same chemical compositions, concerns exist about their potential human health effects. In particular with regard to ceramic nanofibres, there is a paucity of information concerning risk from inhalation exposure.

The effect of fibre characteristics on lung deposition, clearance, and pulmonary response should be systematically evaluated to establish the true biological effect of high aspect ratio nanomaterials. ^{9–12} For micron-scale fibres, the accepted paradigm is that toxicity is determined by the fibre dose, dimensions, and durability. ^{13,14} The lung dose delivered over time is determined by fibre deposition (respirability). ^{15–17}

The dominant factor that controls fibre respirability is the aerodynamic diameter, which controls the transport and deposition efficiency of the fibres in different regions of the lung. ¹⁷ The fibre durability or biopersistence is the retention of fibres in the lung, over time, with regard to number, dimension, surface area, chemical composition, and similar physical characteristics. ¹⁵

A fibre is considered biopersistent if the rate at which it dissolves *via* chemical dissolution is slow relative to the rate of physical removal from the lung by mechanical action (alveolar macrophage and mucociliary-mediated clearance). ¹⁸ In the non-ciliated alveolar (gas exchange) region of the lung, a primary mechanism for clearance of fibres is engulfment by alveolar macrophages (AM) and movement of the fibre-laden cells to the ciliated bronchioles. ¹⁹ For the AM to be effective in removing fibres, the fibres should be completely engulfed by the AM. If the fibres are too long, the process is not effective or frustrated. Short fibres, less than about 15 to 20 µm in length, are more easily cleared from the lungs by the AM and mucociliary transport than longer fibres. ¹⁴ Long fibres can be removed if they dissolve and break into small fibres. Transverse breakage transforms long fibres into short ones, which may be cleared by AMs and reduce their biological reactivity. Long fibres are more difficult to remove from the lung and are of primary health concern. Biopersistent fibres that remain in the alveoli for an extended period of time can cause harmful effects in humans. ^{14,16,17,19}

Recent studies show that physicochemical characteristics of engineered micron and nanomaterials, (size, shape, surface charge, surface area, and composition) can play significant roles in determining their biological response. ^{20–25} For example, cubic particles with 2 µm side lengths and cylindrical particles of 1 µm length were internalized by a large fraction of cells, whereas cubic particles with side lengths of 3 µm and 5 µm were not taken up to any appreciable degree. ²¹ The shapes of TiO₂ nanomaterials were found to influence toxicity, with elongated structures (fibres) having more biological reactivity than short

structures or spherical particles of the same chemical composition. 23 ZnO and TiO₂ materials with of different shapes, sizes, surface areas, and crystal structures had different cytotoxicity with human lung epithelium cells. 24

The concern of fibre persistence in the lung motivates this work to measure the rate of dissolution of the alumina (Al_2O_3) sub-micron sized fibres. Alumina-based fibres are partially soluble in lung fluids and may not cause pulmonary inflammation,²⁶ which makes them a candidate for development of high aspect ratio nanomaterials that are safe by design.

Unfortunately, little information exists in regard to the physicochemical effects of submicron alumina fibres, which include different fibre sizes, surface morphologies, crystal structures, and surface areas with respect to alumina fibres formed by calcination of electrospun polymeric fibres. The aim of this work is to fill in part of the knowledge gap by resolving the influence of these physicochemical properties on the rates that alumina fibres dissolve in artificial lung fluids and evaluate their potential to generate free radicals.

Experimental

Materials

An aluminium precursor solution was prepared by combining formic acid, aluminium acetate (basic hydrate, $(CH_3CO_2)_2AIOH \cdot_X H_2O$, Alfa Aesar), and distilled water at a weight ratio of 1:1:2.5 respectively. This mixture was well stirred. A 9 wt% polymer solution was prepared by dissolving polyvinylpyrrolidone (PVP, Aldrich, MW: 1 300 000) in ethanol (AAPER alcohol, 200 proof). The electrospinning solution was prepared by a 1:1 mass ratio of the polymer solution with the aluminium precursor solution for 1 hour at room temperature until completely blended, as indicated when the initially cloudy solution turned clear.

Fabrication of aluminium acetate/PVP composite sub-micron fibres

A porous hollow tube multi-jet electrospinning technique was used to fabricate aluminium acetate/PVP composite fibres.²⁷ The multiple jet device was composed of a 1/2 inch PVC pipe with holes, an electrode, a power supply, a grounded collector, and a reservoir as shown in Fig. 1. The electrode was inserted into the PVC pipe to induce an electric charge into a solution.

The electrospinning solution was loaded into the reservoir from which the solution flowed to the pipe. Multiple pendent drops were formed at the holes of the pipe. A voltage potential of about 30 kV DC, high enough to overcome the interfacial tension of the drops, ²⁸ was applied to the solution in the pipe to launch the jets towards a grounded collector. ^{1–5,28}

Thermal treatment of electrospun Al₂O₃ fibres

The electrospun polymer fibres were calcined in air to convert the polymer to ceramic form. The PVP polymer degrades at temperatures higher than 450 °C. In this work, the fibres were calcined using different controlled temperature heating cycles to obtain different crystalline structures: (A) rapid heating to varying soak temperatures (650, 750, 850, 950, 1050, and

1150 °C), and (B) different ramp rates (93, 204, 315, and 600 °C h⁻¹) to the same soak temperature of 650 °C. The soak times were the same for all temperature cycles (4 h).

After heating, the samples were allowed to cool slowly in the furnace. The temperature profiles for the various cycles are shown in Fig. 2. The fibre samples labelled as S650, S750, S850...S1150 correspond to the different soak temperatures and samples labelled as R93, R204, R315, and R600 correspond to the different ramp rates. As a control, the sample labelled PVP01 corresponds to uncalcined aluminium precursor – PVP fibres. The characterization and dissolution results of the fibre samples are referenced to these labels.

Physical characterization of materials

A high-resolution transmission electron microscope (HR-TEM, FEI Technai F30) operating at 300 kV and a Transmission Electron Microscope (TEM, JEM 1200XII) were used to study fibre size and morphology of the Al_2O_3 fibres. Small samples of the fibre mats were cut and placed in acetone, and ultrasonically vibrated to form suspensions. Drops of the suspensions were passed through a carbon coated copper grid, leaving the fibres on the grid for TEM imaging.

Scanning electron microscope (SEM, FEI Quanta 200 at 30 kV and HITACHI TM300 at 15 kV) images were analysed to study the fibre morphology and determine fibre size distributions. The diameter distributions were determined from more than 100 fibre diameter measurements for each sample from the SEM images using FibraQuant 1.3 software (nano-Scaffold Technologies LLC).

An X-ray diffractometer (Rigaku D/Max-2000T) with Cu anode (K α_1 = 0.154056 nm) was used for determination of the crystal phase, crystallinity, and crystal size of the Al₂O₃ fibres. The voltage was set to 40 kV and the current was set to 40 mA. Scans were collected with a step size of 0.1 deg and scan speed of 0.5 deg per min. The crystallite size of alumina fibres was calculated using the Scherrer equation, which relates the full width at half maxima (FWHM), W, of the most intense peak (440) of γ -Al₂O₃ fibre to the angle of incidence, θ , $via\ S = c\lambda/W\cos\theta$, where S is the average crystallite size, c is the Scherrer constant and λ is the wavelength. The Scherrer constant can take values from 0.9 to 1.2, depending on the shape of the particles. Here we assumed the spherical shape (c = 1).

A Fourier Transform Infrared Spectrometer (Thermo Fisher Scientific Model: Nicolet iS50 FT-IR) was used to determine the molecular bonding information of the samples. The samples were prepared by drying at 75 °C for 2 days to completely remove the moisture, and each sample was pressed into discs for the tests. The IR spectra were recorded from 400 to 4000 cm^{-1} at 8 cm^{-1} resolution.

The Brunauer, Emmett and Teller (BET) surface areas were measured using nitrogen gas adsorption (ASAP2020, Micromeritics Corp., Norcross, GA). A value of 0.162 nm² was used for the molecular cross-sectional area of N_2 at 77 K. The BET was calculated from at least six adsorption points in the range $p/p_0 = 0.01$ to 0.3; the values were normalized by dry sample mass to calculate specific surface area with units of m² g⁻¹.

Powder density was measured using a helium pycnometer (AccuPycII 1340, Micromeritics) using research grade gas at 15.0 psig. Prior to the density measurements, the fibre samples were placed in an oven at 100 °C to dry overnight and allowed to cool to ambient temperature in a desiccator. A total of 25 consecutive volume measurements were made with the equilibration rate set at 0.0005 psig min⁻¹ and the results normalized by dry sample mass to calculate density with units of g cm⁻³.

Artificial lung fluid dissolution studies

The biodurability of the Al_2O_3 fibres was assessed in two different artificial lung fluid models viz. lung airway epithelial lining fluid having near-neutral pH²⁹ and alveolar macrophage phagolysosomal fluid having acidic pH.³⁰ Material dissolution was evaluated using the well-established static technique.³¹ With this technique, a known mass of each fibre sample was weighed onto a 0.025 μ m pore size 47 mm diameter nitrocellulose filter (Millipore, Bedford, MA) and a second filter of the same type was placed on top to create a 'sandwich'. The filters served as barriers to particle migration into the dissolution medium which would bias estimates of dissolved alumina. Each filter sandwich was clamped in a dissolution chamber assembly and secured using an O-ring and Teflon screws (Fig. 3) Importantly, the filter pore size (25 nm) was significantly smaller than the diameter of the fibres.

The fraction of each fibre sample that was aluminium metal was determined by instrumental neutron activation analysis (Elemental Analysis Inc., Lexington, KY) and used to calculate the initial mass of aluminium from the gravimetrically determined fibre masses. Each filter sandwich was secured tightly in a static dissolution chamber (Intox Products, Moriarity, nm) using four nylon screws and O-rings to provide a particle-tight seal around the edge of the sandwich. Each dissolution chamber assembly was placed in a separate plastic container with 0.080 L of lung fluid and the masses of dissolved aluminium were quantified at predesignated time points. Triplicate dissolution chambers were prepared for each Al₂O₃ fibre. To mimic human lung temperature all samples were maintained at 37 °C for the duration of the respective study periods using a water-jacketed incubator.

To model the fate of particles that deposit in conducting airways, biodurability was evaluated in artificial lung airway epithelial lining fluid termed serum ultrafiltrate (SUF). Generally, particles that deposit in the conducting airways are bathed in the epithelial lining fluid and cleared by mucociliary action within several hours. Hence, samples of SUF were quantified for dissolved aluminium at 3 h, 6 h, 12 h, 24 h, 2 day, 4 day, and 7 day (the longer time points were collected to accurately capture the fibre dissolution kinetics). The pH of SUF was maintained at 7.3 ± 0.1 through-out the study by passing flowing CO_2 (5%) and air (95%) across the solvent.

To model the fate of particles that deposit in the alveoli, biodurability was evaluated using macrophage phagolysosomal simulant fluid (PSF).³⁰ Small particles with aerodynamic diameters below 4 µm can deposit in the distal non-ciliated alveolar (gas exchange) region of the lung. Upon deposition in the alveoli, particles are quickly engulfed by scavenger cells such as macrophages and isolated in vacuoles called lysosomes. The lysosomes merge with phagosomes that contain an acidic milieu to form phagolysosomes that function to sequester

and degrade foreign materials over days to weeks. To understand dissolution kinetics over the lifespan of a macrophage cell, masses of dissolved aluminium were quantified following particle immersion for 3 h, 6 h, 12 h, 24 h, 2 day, 4 day, 7 day, 14 day, 21 day, and 28 day. The pH of PSF was maintained at 4.5 ± 0.1 using 0.02 M potassium hydrogen phthalate buffer.

Analysis of dissolved aluminium was performed using inductively coupled plasma-optical emission spectroscopy without sample digestion. The analytical limit of detection (LOD) for aluminium in SUF was 5 μ g L⁻¹ and the limit of quantification (LOQ) was 17 μ g L⁻¹. For PSF, the aluminium LOD was 10 μ g L⁻¹ and the LOQ was 43 μ g L⁻¹.

Dissolution rates of the fibres were calculated assuming constant dissolution velocity³²:

$$k = \frac{2 \cdot \left(1 - \frac{M}{M_0}\right)^{\frac{1}{2}}}{\text{SSA} \cdot t} \quad (1)$$

where $(1-M/M_0)$ is the mass fraction of material dissolved, t is time (days), SSA is the specific surface area (cm² g⁻¹), and k is the chemical dissolution rate constant (g per cm² per day). For each material, values of $(1-M/M_0)$ were potted *versus* t, and k was calculated from the slope (b) of the plot for $t \to 0$ and normalized to SSA/2. The dissolution half-time ($t_{1/2}$) calculated as 0.693/b. In general, the dissolution was biphasic and consisted of an initial rapid phase followed by a slower long-term phase. Hence, curves were fitted to the initial and long-term phases separately using a non-linear least squares fitting procedure (SAS 9.2, SAS Institute Inc., Cary, NC). Analysis of variance (ANOVA) F-statistics were used to note overall differences in $t_{1/2}$ and k values among materials. Tukey's test was used to identify specific paired differences between means.

Free radical generation studies

Macrophages attack and engulf foreign bodies (including particles) that invade the lung where they can react with H_2O_2 as one of their defence mechanisms. This H_2O_2 can react with particles in a Fenton-like manner and generate more damaging free radicals. Modelling this respiratory burst defence system with H_2O_2 provides a source of preliminary data on the biological reactivity of the particles involved.³³ Hence the Al_2O_3 fibres were reacted with H_2O_2 to measure their potential to generate reactive radical species when inhaled and exposed to an organism's defence systems. Hydroxyl radical production(OH–) was measured using the addition-type reaction of a short-lived radical with a compound (spin trap) to form a relatively long-lived free radical product (spin adduct), which can be studied using conventional electron spin resonance (ESR) spin trapping.

For H_2O_2 Fenton-like reaction measurements, reactants were mixed in test tubes at a final volume of 1.0 mL of phosphate buffered saline (PBS), Al_2O_3 fibres in the presence of 10 mM H_2O_2 and 100 mM of 5,5-dimethyl-1-pyrroline-*N*-oxide (DMPO) (final fibre concentration of 1 mg mL⁻¹) and vortexed for 10 seconds. In this system, a high dose was required to observe a reaction within the short exposure time necessitated by the use of ESR. The reaction mixture was transferred to a flat cell for ESR measurement (Bruker EMX spectrometer, Bruker Instruments Inc., Billerica, MA) at room temperature under ambient

air. Hyperfine couplings were measured (to 0.1 G) directly from magnetic field separation using potassium tetraperoxochromate and 1,1-diphenyl-2-picrylhydrazyl as reference standards. 34,35 Min–U–Sil silica (<5 μ m) was used as a particle control. The relative radical concentration was estimated by multiplying half of the peak height by (H_{pp}), where H_{pp} represents peak-to-peak width. The Acquisit program (Bruker Instruments Inc.) was used for data acquisitions and analyses.

Results and discussion

Al₂O₃ fibre characterization

The unique aspect of our study design was that the correlations between dissolution and physicochemical properties of Al_2O_3 fibres were determined by changes in the fibre size, crystallite size, phase transformation, crystallinity, and specific surface area. These changes are important for understanding nanomaterial dissolution behaviour. Depending on the various characteristics of alumina fibres from the variations in the thermal treatments, the dissolution rates and toxicity will change.

SEM, TEM, and fibre size distribution

For the analyses of the Al_2O_3 fibres calcined at soak temperatures 650 °C, 750 °C, 850 °C, and 950 °C for 4 h, the diameters of Al_2O_3 fibres gradually decreased with increase in the calcination temperature as indicated in Fig. 4 and 5 shows the different ramp rates did not have a significant effect on the fibre diameters.

All of the calcined fibres had diameters smaller than the precalcined PVP polymer fibres due to evaporation of residual solvent and the degradation of the polymer. SEM and TEM images and the corresponding fibre diameter frequency distributions of the uncalcined fibres and the fibres calcined at different temperatures are shown in Fig. 4.

Each distribution of the fibre diameters was measured from at least 10 SEM images. Each histogram was generated out of 500 measurements. It is evident from the distribution plots that the distribution became narrower with calcinations at higher temperature. The S950 samples had the lowest average fibre diameter and the narrowest diameter distribution among all the materials. This trend showed that different calcination temperatures directly affected the diameters of the fibres, and is consistent with observations reported in literature.⁴

The TEM images in Fig. 4 show the fibres with higher calcination temperatures appear to have produced a rougher surface morphology. The apparent roughness corresponds to the increase in crystallinity of the alumina at the higher temperatures. The changes in surface structural features can be attributed to a difference in crystal formation and transformations from amorphous to γ -phase of alumina fibres (described below).

The SEM, TEM and fibre diameter distributions for R93–R600 are shown in Fig. 5. Increasing the temperature ramp rate from 93 $^{\circ}$ C per hour to 600 $^{\circ}$ C per hour appears to not have significantly changed the amorphous nature of the fibres. The smooth fibre surface shown in Fig. 5(B) indicated that there was not an obvious phase change to a crystalline

structure among these samples (R93–R600) with different heating rate. The average fibre diameters were about the same and the distributions had about the same standard deviations.

XRD characterization

To confirm the change of the crystal structure with calcination and to correlate with the observed microscopy data, X-ray diffraction analyses were undertaken. Fig. 6 shows no crystalline peak on the samples before calcinations (PVP01) and the fibre samples calcined at soak temperature 650 °C (*i.e.* samples S650 and the fibres calcined at the various ramp rates R93–R600).

The XRD patterns of samples with calcination temperatures of 750 °C and greater (S750–S950) showed γ -phase peaks at 2θ = 32.8°, 37.6°, 39.6°, 45.9°, 61.1°, and 67.2° representing the indices of (220), (110), (222), (400), (511), and (440) crystal planes of Al₂O₃. The crystallinity of the samples increased with the soak temperature from 13.2% to 40.6% with 8 nm crystallite size of γ -Al₂O₃ fibres as displayed in Fig. 6.

FT-IR characterization

The evolution of the IR spectra of the alumina fibres with different heating conditions are shown in Fig. 7. Comparison of the precalcined PVP polymer fibre (PVP01) with the calcined fibres in Fig. 7(A) shows all of the peaks associated with the polymer aluminium acetate/PVP composite fibres (PVP01) disappeared from the spectra of the Al_2O_3 fibres (S650–S950). The broad and smooth absorption bands in the ranges of 400–1100 cm⁻¹, related respectively to the Al–O (AlO₆ and AlO₄) and O–H stretching vibration, reveal the formation of the amorphous structure of the Al_2O_3 fibres, indicated in Fig. 7(B) and (C). The broad band spectra of the γ -Al₂O₃ observed in Fig. 7(B) is similar to that reported in literature.^{36–39} The absorption band of γ -Al₂O₃ in the range 500 to 1100 cm⁻¹ could be related to the stretching vibration of the whole spinel unit cell.³⁷

The broad absorption at $1048~cm^{-1}$ of the samples heated at 650–950~°C indicates Al–O bond formation. With increasing calcination temperature, there was no significant increase in the intensity of the peaks. Fig. 7(C) shows that the symmetric stretching vibration AlO₄ bands were located at 713 and 771 cm⁻¹. These two stretching bands depend strongly on the crystallinity of γ -Al₂O₃ phase formation. Basically, the cation vacancy distribution leads to the appearance of the broad band due to disordered vacancies. It means that the formation of peaks at 713 and 771 cm⁻¹ are associated with the larger lattice distortion. 37,41 Comparing intensity of the adsorption bands of samples S750, S850 and S950 in Fig. 7(C), shows the γ -Al₂O₃ was formed with increasing calcinations temperature.

BET, INAA and helium pycnometry characterization

Fig. 8 shows the specific surface areas for all of the samples. The specific surface areas increased up to soak temperature 850 °C followed by a decrease in area at soak temperature 950 °C. To determine if the decline in surface area continued at higher temperatures, additional samples were calcined at temperatures 1050 and 1150 °C (S1050 and S1150). The surface areas of the additional samples plateaued at about 16 g m⁻². Samples calcined at the different ramp rates to the same soak temperature (R93–R600) did not show significant

change in the specific surface area, indicating the soak temperature is the more important factor.

BET and XRD characterization were performed to comprehend the correlation between the reduction of surface area and crystal structure. Generally, during the transition from γ to α - Al_2O_3 crystal phases, a dramatic reduction in surface area occurs. 42,43 Upon heating, the γ - Al_2O_3 goes through a series of transformations to reach the thermodynamically stable α - Al_2O_3 phase structure. 42,44,45 Thus, the surface grain coarsening of the Al_2O_3 fibre incurred during calcination can lead to reduction of the specific surface area. 44

Fig. 9 shows that alumina fibres calcined at 1050 °C (S1050) and 1150 °C (S1150) had alpha crystal structures with 15 nm and 17 nm of estimated crystallite sizes. The increase in peak intensities with increase in temperature shows the progressive increase in the alpha phase content of the fibres and the coalescence of the tiny crystallites started accelerating the grain growth to go to alpha phase. From XRD and TEM studies, crystallite growth was also demonstrated by the increase of calcinations temperature, which may reveal how phase transformation modified the fibre morphology and crystal structure.

Fig. 10 shows the atomic alumina content and fibre intrinsic density of the fibres. Within the error of the measurements (relative standard deviation = 6%), the Al content of S950, R204, and R315 matched the value expected from stoichiometry of Al_2O_3 (52.9%). The calcination temperature clearly made a difference in Al content from PVP01 to S950 in Fig. 10(top). However, regardless of ramp rate, samples R93 to R315 had similar (but slightly less than theoretical) Al content apart from R600 samples. Al content of R600 was approximately 40%. In Fig. 10(bottom), the density of the samples also increased with calcination temperature, which was also related to agglomerated grains on the surface of the alumina fibres calcined at high temperature. Density of R93, R204...R600 samples having different ramping rate had no significant change of the values.

Artificial lung fluid dissolution studies

On a mass basis, the total amounts of aluminium dissolved from the fibres ranged from 0.2 to 13.5% in SUF (Table 1) and from 0.5 to 36% in PSF (Table 2). In both lung fluid models, the largest total masses of dissolved aluminium were from the uncalcined PVP01 material, which reflects the greater solubility of the precursor aluminium acetate relative to the heat-treated alumina fibres.

Aluminium dissolution was biphasic in both lung fluid models. Estimates of the dissolution half-time, $t_{1/2}$ (the time to clear half the mass via solubilization) and the chemical dissolution rate constant (k) for the initial rapid phase and latter long-term phase of each Al₂O₃ fibre are summarized in Table 1 (SUF) and in Table 2 (PSF). In both SUF and PSF, k values for PVP01 were significantly faster than all of the alumina fibres (p < 0.05).

Inhaled alumina fibres that deposit in the conducting airways will be subject to clearance *via* chemical dissolution and mechanical action (mucociliary-mediated transport). Particle clearance by chemical dissolution depends upon the composition of the airway lining fluid, chemistry of the particle, *etc*. In contrast, mechanical clearance is independent of these

factors and transport to the pharynx (where the particles are swallowed and excreted *via* the gastrointestinal tract) occurs at rapid rates with half-times of approximately 2 to 8 h.⁴⁶

Hence, for SUF, the calculated $t_{1/2}$ and k values indicate that if Al_2O_3 fibres deposit in the conducting airways of the lung, little material will dissolve prior to removal by mechanical action.

Once deposited in the non-ciliated alveolar (gas exchange) region of the lung, particles are rapidly engulfed by AM and sequestered within phagolysosomes. The engulfed particles are cleared *via* chemical dissolution in the acidic microenvironment of the phagolysosomes and by movement of particle-laden cells to the ciliated bronchioles, ¹⁹ and eventually to the pharynx. Movement of AM to the ciliated bronchioles is a very slow process with half-times of approximately 35 to 7000 days. ⁴⁶

Generally, the initial and latter phase dissolution half-times of the alumina fibres in PSF were less than 7000 days. The greater solubility of alumina fibres in PSF relative to macrophage-mediated clearance indicates that chemical dissolution is an important clearance mechanism if these fibres were to deposit in the alveoli.

The exception to this observation was the latter-phase dissolution half-times for R93, R204, and R315 which were a factor of two longer than expected for mechanical clearance and would be expected to have prolonged persistence in the alveoli. The slow dissolution and potential persistence of aluminium oxide fibres in the lung alveoli has important implications for understanding human health. Released ions can bind to proteins such as albumin and/or be eliminated through urine. Persistence of particulate can induce pulmonary damage as a consequence of accumulation of non-reactive particles. ⁴⁷ Aluminium is ubiquitous in our environment and is present in trace amounts in the human body, as such, inhalation and dissolution of aluminium particulate could lead to biomagnification of levels in the body and alter homeostasis.

There were no statistical differences in the initial rapid or latter long-term dissolution rates among S650 (650 °C), S750 (750 °C), S850 (850 °C), and S950 (950 °C) in either SUF or PSF. Hence, our data indicate that dissolution rates were independent of calcination temperature. Next, we evaluated the influence of calcination ramp rate on dissolution using the fibres that were calcined at the same temperature (650 °C) but using heating ramp rates of 93 (R93), 204 (R204), 315 (R315) or 600 (R600) °C per hour.

The dissolution clearance half-time of R600 was significantly shorter compared to R93, R204, and R315 (p < 0.05). The long term dissolution rates of aluminium from R600 were a factor of five faster in SUF and a factor of 14 faster in PSF relative to R93–R315, though differences were not significant. These data suggest an effect of calcination ramp rate on dissolution (all four types had similar diameters and densities and were amorphous). Specifically, a rapid ramp rate (600 °C per hour) to 650 °C yielded a less persistent alumina fibre. For a metallurgical study regarding the effect of ramp rate, some authors showed that densification and fine grain sizes of alumina materials could be changed. $^{48-50}$

There have been conflicting results. Stanciu *et al.* and Zhou *et al.* reported that the grain size of alumina decreased with increasing heating rate under 45 MPa pressure and high densities was obtained in alumina ceramic materials without any additives. ^{48,49} Whereas, Kim *et al.* stated that the grain size and residual porosity decreased with slow heating rate. ⁵⁰ Hence, heating ramp rate might affect dissolution results due to the change of the densification and the direct grain to grain contact at molecular scale in alumina ceramic fibre.

Previously, Adamcakova-Dodd *et al.* evaluated the dissolution of a commercially available Al_2O_3 fibre material using a model of airway epithelial lining fluid and a model of macrophage lysosomal fluid analogous to SUF and PSF, respectively. Though sold as Al_2O_3 powder, characterization of the as-received material revealed that it was primarily composed of crystalline aluminium hydroxide. The authors reported that the dissolution of aluminium hydroxide was generally linear in both lung fluid models. From their data, we estimated that $k = 9 \times 10^{-10}$ g per cm² per day in their airway lining fluid model and $k = 4 \times 10^{-9}$ g per cm² per day in their macrophage lysosomal fluid. Our k value estimates for the long-term dissolution of amorphous (with the exception of PVP01, S650, and R600) and crystalline alumina fibres were similar to that derived for aluminium hydroxide.

Visual inspection of Al₂O₃ fibres after 28 days of exposure to PSF indicated that these materials were susceptible to transverse breakage (Fig. 11). For macrophages to be effective in removing fibres from the alveoli, the fibres should be completely engulfed by the cells. If the fibres are too long, mechanical clearance *via* particle-laden macrophage migration from the alveoli is less effective. Our observation that the Al₂O₃ fibres break transversely from exposure to PSF is important because breakage transforms long fibres into shorter fibres which are more easily cleared by AMs *via* migration from the alveoli.¹³

Free radical generation

None of the Al₂O₃ fibres generated measureable levels of 'OH radicals in the cell-free H₂O₂ test system. Both of the soluble hexavalent chromium and particulate silica positive control samples generated high levels of hydroxyl radicals. The absence of radical production by these Al₂O₃ fibres is consistent with Adamcakova-Dodd *et al.* who reported that inhalation of aluminium hydroxide whiskers by mice did not produce lung inflammation.²⁶

It is important to note that macrophage engulfment of particles and reaction with ${\rm H_2O_2}$ was screened using a cell-free system. In cellular systems, particle size, shape, and/or surface charge influence uptake by cells. $^{20-23}$ For example, titanium dioxide nanobelts having length of <5 μ m are fully taken up into cell lysosomes whereas longer nanobelts with length >15 μ m are only partially internalized with the rest sticking out into the cytoplasm. 23 Depending on their degree of internalization and interaction with organelles, particulate can promote various biological responses, including cell proliferation, free radical generation, release of pro-inflammatory cytokines, cytotoxicity, apoptosis, cytoskeleton formation, and adhesion and migration.

Conclusions

The present study concluded the biodurability of eight electrospun alumina fibre modified by different heating protocol in artificial lung fluid models. We found that first, uncalcined PVP-Al₂O₃ fibre (PVP01) was significantly faster than all other alumina fibres, and alumina fibres calcined from 650 °C to 950 °C (S650–S950) did not cause the change of dissolution half-time significantly. Thus, the physicochemical property of Al₂O₃ fibre had no effect on dissolution behaviour in either SUF or PSF. For the variation in heating ramp rate (R93–R600), the alumina fibres from R600 had a significantly greater dissolution rate compared to slowly heated alumina fibre samples (R93–R315) even though these four samples had similar physicochemical property such as, fibre diameter, crystal structure (amorphous), surface area, and density. The fibres with increased solubility have a reduced inflammogenic potential. To fully understand the effect of the ramping rate, additional work is needed.

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Notes and references

- 1. Lin, T. Nanofibres: Production, properties and functional applications. InTech; 2011.
- 2. Ramaseshan R, Sundarrajan S, Jose R, Ramakrishna S. J Appl Phys. 2007; 102:111101–111117.
- 3. Shin HU, Li Y, Paynter A, Nartetamrongutt K, Chase GG. Polymer. 2015; 65:26–33.
- 4. Park SJ, Bhargava S, Bender ET, Chase GG, Ramsier RD. J Mater Res. 2008; 23:1193-1196.
- 5. Barhate RS, Loong CK, Ramakrishna S. J Membr Sci. 2006; 283:209-218.
- 6. Brown, RC. Air filtration: an integrated approach to the theory and applications of fibrous filters. Pergamon Press; 1993.
- 7. Graham, K.; Ouyang, M.; Raether, T.; Grafe, T.; McDonald, B.; Knauf, P. Fifteenth Annual Technical Conference & Expo of the American Filtration & Separations Society; 2002. p. 9-12.
- 8. National Research Council of the National Academies. A Research Strategy for Environmental, Health, and Safety Aspects of Engineered Nanomaterials. The National Academies Press; 2012.
- 9. Misra SK, Dybowska A, Berhanu D, Luoma SN, Valsami-Jones E. Sci Total Environ. 2012; 438:225–232. [PubMed: 23000548]
- 10. Braakhuis H, Park M, Gosens I, De Jong W, Cassee F. Part Fibre Toxicol. 2014; 11(18):1–25. [PubMed: 24382024]
- 11. Borm P, Klaessig FC, Landry TD, Moudgil B, Pauluhn J, Thomas K, Trottier R, Wood S. Toxicol Sci. 2006; 90:23–32. [PubMed: 16396841]
- 12. Landsiedel R, Ma-Hock L, Kroll A, Hahn D, Schnekenburger J, Wiench K, Wohlleben W. Adv Mater. 2010; 22:2601–2627. [PubMed: 20512811]
- 13. Donaldson K, Murphy F, Schinwald A, Duffin R, Poland CA. Nanomedicine. 2010; 6:143–156. [PubMed: 21182425]
- 14. Davis JMG. Environ Health Perspect. 1994; 102:113–117. [PubMed: 7882914]
- 15. McClellan RO, Hesterberg TW. Environ Health Perspect. 1994; 102:277–283. [PubMed: 7882950]
- 16. Hesterberg TW, Hart GA. Crit Rev Toxicol. 2001; 31:1–53. [PubMed: 11215691]

17. Maxim LD, Hadley JG, Potter RM, Niebo R. Regul Toxicol Pharmacol. 2006; 46:42–62. [PubMed: 16837114]

- 18. Stefaniak A. Part Fibre Toxicol. 2010; 7(38):1–9. [PubMed: 20180970]
- Bernstein D, Castranova V, Donaldson K, Fubini B, Hadley J, Hesterberg T, Kane A, Lai D, McConnell EE, Muhle H, Oberdorster G, Olin S, Warheit DB. Inhalation Toxicol. 2005; 17:497– 537.
- 20. Misra SK, Nuseibeh S, Dybowska A, Berhanu D, Tetley TD, Valsami-Jones E. Nanotoxicology. 2014; 8:422–432. [PubMed: 23590525]
- 21. Gratton SEA, Ropp PA, Pohlhaus PD, Luft JC, Madden VJ, Napier ME, DeSimone JM. Proc Natl Acad Sci U S A. 2008; 105:11613–11618. [PubMed: 18697944]
- 22. Huang X, Teng X, Chen D, Tang F, He J. Biomaterials. 2010; 31:438–448. [PubMed: 19800115]
- 23. Hamilton R, Wu N, Porter D, Buford M, Wolfarth M, Holian A. Part Fibre Toxicol. 2009; 6(35):1–11. [PubMed: 19134195]
- 24. Hsiao IL, Huang YJ. Sci Total Environ. 2011; 409:1219–1228. [PubMed: 21255821]
- 25. Mu Q, David CA, Galceran J, Rey-Castro C, Krzemi ski Ł, Wallace R, Bamiduro F, Milne SJ, Hondow NS, Brydson R, Vizcay-Barrena G, Routledge MN, Jeuken LJC, Brown AP. Chem Res Toxicol. 2014; 27:558–567. [PubMed: 24575710]
- 26. Adamcakova-Dodd A, Stebounova L, O'Shaughnessy P, Kim J, Grassian V, Thorne P. Part Fibre Toxicol. 2012; 9(22):1–14. [PubMed: 22239852]
- 27. Varabhas JS, Chase GG, Reneker DH. Polymer. 2008; 49:4226-4229.
- 28. Reneker DH, Yarin AL, Fong H, Koombhongse S. J Appl Phys. 2000; 87:4531-4547.
- Finch GL, Mewhinney JA, Eidson AF, Hoover MD, Rothenberg SJ. J Aerosol Sci. 1988; 19:333–342.
- 30. Stefaniak AB, Guilmette RA, Day GA, Hoover MD, Breysse PN, Scripsick RC. Toxicol in vitro. 2005; 19:123–134. [PubMed: 15582363]
- 31. Kanapilly GM, Raabe OG, Goh CHT, Chimenti RA. Health Phys. 1973; 24:497–507. [PubMed: 4707664]
- 32. Potter RM, Mattson SM. Glastech Ber. 1991; 64:16–28.
- 33. Dahlgren, C.; Karlsson, A.; Bylund, J. Neutrophil Methods and Protocols. Quinn, M.; DeLeo, F.; Bokoch, G., editors. Vol. 412. Humana Press; 2007. p. 349-363.ch. 23
- 34. Janzen EG, Blackburn BJ. J Am Chem Soc. 1968; 90:5909-5910.
- 35. Buettner GR. Free Radical Biol Med. 1987; 3:259–303. [PubMed: 2826304]
- 36. Tarte P. Spectrochim Acta, Part A. 1967; 23:2127-2143.
- 37. Saniger J. Mater Lett. 1995; 22:109-113.
- 38. Pecharromán C, Sobrados I, Iglesias JE, González-Carreño T, Sanz J. J Phys Chem B. 1999; 103:6160–6170.
- 39. Du X, Wang Y, Su X, Li J. Powder Technol. 2009; 192:40-46.
- 40. Meher T, Basu AK, Ghatak S. Ceram Int. 2005; 31:831-838.
- 41. Mo C, Yuan Z, Zhang L, Xie C. Nanostruct Mater. 1993; 2:47–54.
- 42. Das RN, Hattori A, Okada K. Appl Catal, A. 2001; 207:95-102.
- 43. Kwak JH, Peden CHF, Szanyi JN. J Phys Chem B. 2011; 115:12575–12579.
- 44. Levin I, Brandon D. J Am Ceram Soc. 1998; 81:1995-2012.
- 45. Yu P-C, Yang R-J, Tsai Y-Y, Sigmund W, Yen F-S. J Eur Ceram Soc. 2011; 31:723-731.
- 46. ICRP. Human Respiratory Tract Model for Radiological Protection. ICRP Publication. 1994; 66
- 47. Willhite CC, Karyakina NA, Yokel RA, Yenugadhati N, Wisniewski TM, Arnold IMF, Momoli F, Krewski D. Crit Rev Toxicol. 2014; 44:1–80. [PubMed: 25233067]
- 48. Zhou Y, Hirao K, Yamauchi Y, Kanzaki S. J Eur Ceram Soc. 2004; 24:3465–3470.
- 49. Stanciu LA, Kodash VY, Groza JR. Metall Mater Trans A. 2001; 32:2633-2638.
- 50. Kim BN, Hiraga K, Morita K, Yoshida H. J Eur Ceram Soc. 2009; 29:323-327.

Nano impact

Engineered nanomaterials represent a class of materials with novel properties that make them ideal for specific applications. Among nanomaterials, alumina nanofibres are of high interest for thermal stability in high temperature applications. From a health and safety perspective fibre materials should be evaluated for their potential to cause pulmonary diseases. In this work, we compared the biodurability of different types of alumina nanofibres in artificial lung fluid models and elucidate how the physicochemical properties are controlled by the production process to yield fit-for-purpose nanofibres with varying biodurabilities. Engineering parameters can be manipulated to produce fibres with most favourable biodurabilities within the constraints of the application operating conditions. This work is relevant to commercial industries, public and occupational health researchers, and regulators.

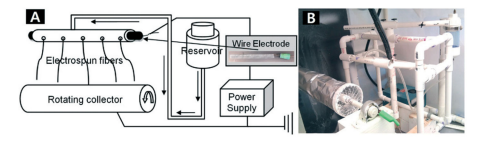


Fig. 1.
Porous hollow tube electrospinning (A) schematic diagram, (B) laboratory setup.

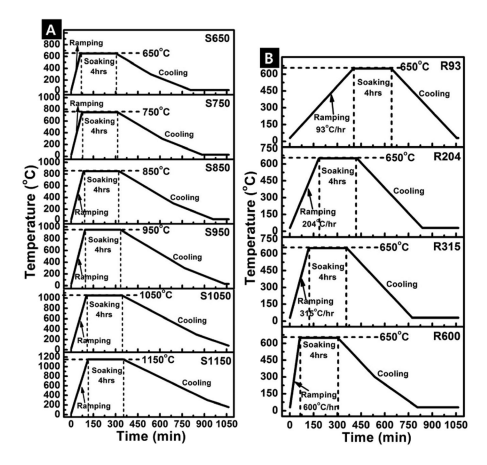


Fig. 2. Temperature profiles of the variations in the heating cycles to calcine the alumina precursor–PVP fibres to fabricate the Al_2O_3 fibres. (A) Variation in soak temperatures (samples labelled S650 to S1150). (B) Variation in heating ramp rates (samples labelled R93, R204, R315, and R600).

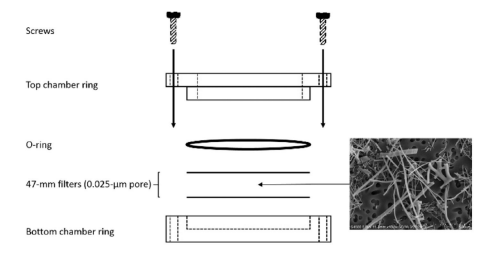


Fig. 3. Static dissolution chamber assembly; SEM micrograph of S950 material after 7 day exposure to SUF.

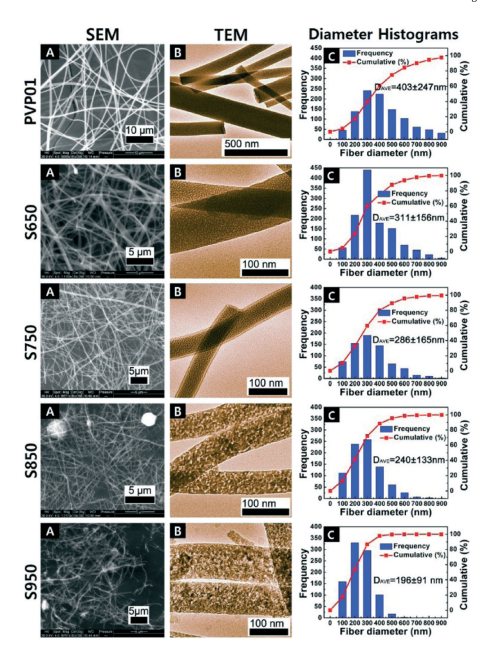


Fig. 4. SEM (A), TEM (B) and length weighted frequency distribution of fibre size; (C) of the uncalcined fibres and the fibres calcined for 4 h at different temperatures.

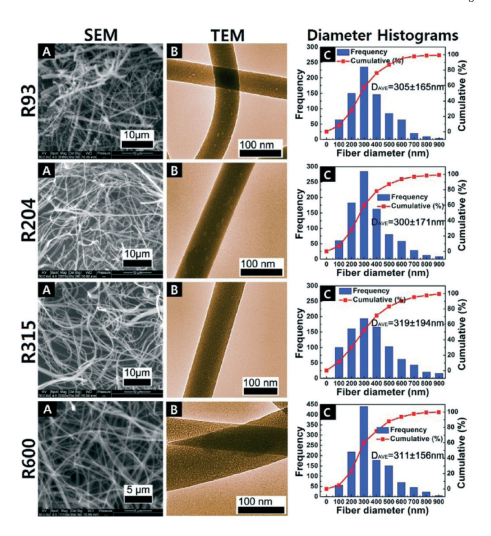


Fig. 5. SEM (A), TEM (B) and fibre diameter distributions (C) of Al_2O_3 fibres calcined at 650 °C for 4 h R93 (93 °C h⁻¹), R204 (204 °C h⁻¹), R316 (316 °C h⁻¹), and R600 (600 °C h⁻¹).

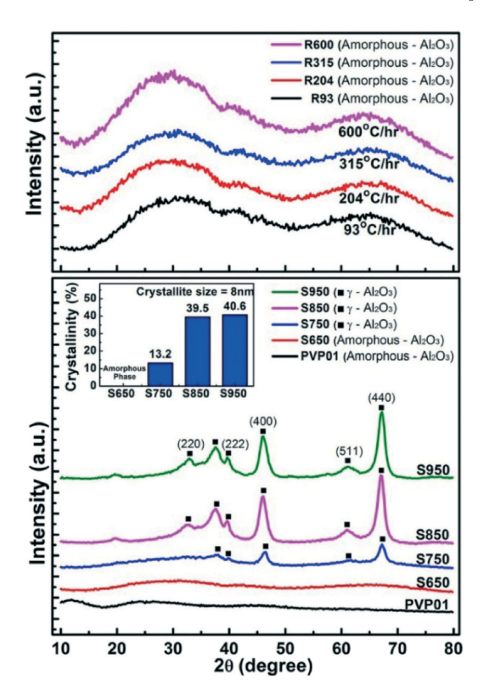


Fig. 6. XRD patterns and crystallinity of Al_2O_3 fibres calcined at different soak temperatures and ramp rates.

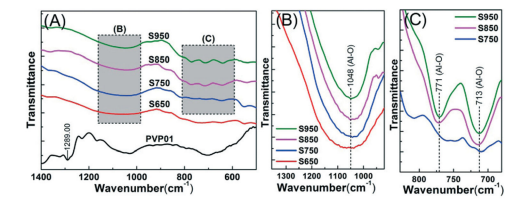


Fig. 7. FT-IR spectra of precalcined polymer PVP composite (PVP01) and the calcined Al_2O_3 fibres at various soak temperatures (S650–S950). (A) Full spectra 500–1400 cm⁻¹; (B) and (C) are reduced spectra ranges as indicated by boxes in (A).

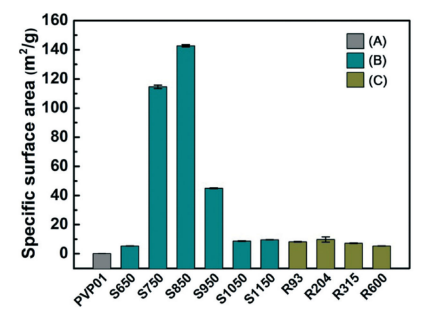


Fig. 8. Specific surface area of alumina fibres. (A) PVP01, (B) fibres rapidly heated to final soak temperatures (S650–S1150); (C) fibres heated at different ramp rates to soak temperature 650 °C (R93–R600).

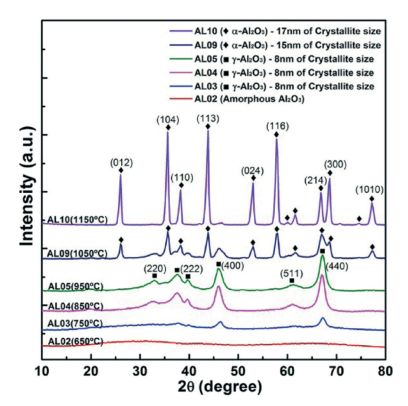


Fig. 9. XRD plots of alumina fibres calcined at 650–1150 °C.

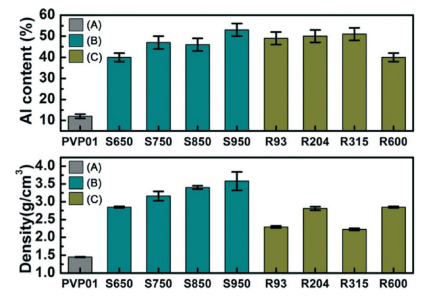


Fig. 10. Al content (top) and density of alumina fibre (bottom) (A) PVP01, (B) S650–S950, and (C) R93–R600.

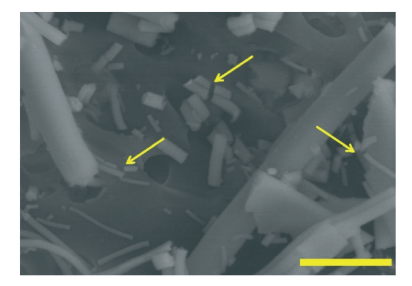


Fig. 11. Scanning electron micrograph of alumina fibre S950 (950 °C) illustrating transverse fibre breakage (see arrows) after exposure to PSF for 28 days. Scalebar = 1 μ m.

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Table 1

Dissolution parameters of alumina fibres in SUF (pH 7.4)

k (g per cm² per day) $1.7 \pm 0.7 \times 10^{-10}$ $1.8 \pm 0.3 \times 10^{-10}$ $1.0 \pm 0.0 \times 10^{-10}$ $4.4\pm2.3\times10^{-6}$ $2.1 \pm 0.5 \times 10^{-9}$ $7.5 \pm 0.8 \times 10^{-9}$ $7.5 \pm 0.8 \times 10^{-9}$ $1.2\pm0.4\times10^{-9}$ $1.4 \pm 0.3 \times 10^{-9}$ Latter long-term phase Dissolution parameters (mean ± standard deviation) 5387 ± 1080 4164 ± 1009 9881 ± 3394 5515 ± 1223 6998 ± 2461 3480 ± 520 1764 ± 185 1764 ± 185 t_{1/2} (days) 125 ± 57 k (g per cm² per day) $4.8 \pm 1.6 \times 10^{-10}$ $2.9 \pm 0.3 \times 10^{-10}$ $3.9 \pm 1.9 \times 10^{-10}$ $1.5 \pm 0.7 \times 10^{-8}$ $5.6 \pm 2.0 \times 10^{-5}$ $2.1 \pm 0.7 \times 10^{-9}$ $1.2 \pm 0.2 \times 10^{-9}$ $1.9 \pm 0.6 \times 10^{-9}$ $1.5 \pm 0.7 \times 10^{-8}$ Initial rapid phase 4933 ± 1880 4761 ± 2573 4058 ± 1553 5874 ± 852 1032 ± 484 1343 ± 385 1689 ± 195 1032 ± 484 t_{1/2} (days) 9 ± 3 Material Dissolved (%) 13.48 ± 6.66 0.29 ± 0.10 1.13 ± 0.07 0.56 ± 0.13 0.36 ± 0.11 0.22 ± 0.07 0.30 ± 0.03 0.49 ± 0.13 1.13 ± 0.07 PVP01 8650aR315 S850 R600 S750 S950 R204 R93

 a S650 is the same as R600.

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Table 2

Dissolution parameters of alumina fibres in PSF (pH 4.5)

k (g per cm² per day) $2.9 \pm 0.5 \times 10^{-10}$ $2.1 \pm 0.1 \times 10^{-10}$ $1.6 \pm 0.6 \times 10^{-10}$ $5.3 \pm 2.0 \times 10^{-10}$ $4.9 \pm 2.0 \times 10^{-10}$ $5.7 \pm 1.4 \times 10^{-10}$ $3.2\pm0.4\times10^{-6}$ $6.8 \pm 1.0 \times 10^{-9}$ $6.8\pm1.0\times10^{-9}$ Latter long-term phase $15\ 609\pm4110$ 16.943 ± 8692 16365 ± 5639 Dissolution parameters (mean ± standard deviation) 3215 ± 1004 5496 ± 961 1958 ± 302 1958 ± 302 2897 ± 201 t_{1/2} (days) 147 ± 18 k (g per cm² per day) $3.9 \pm 0.4 \times 10^{-8}$ $3.7 \pm 0.6 \times 10^{-5}$ $1.8\pm0.6\times10^{-9}$ $2.3 \pm 0.1 \times 10^{-9}$ $1.1 \pm 0.5 \times 10^{-9}$ $2.3 \pm 1.4 \times 10^{-9}$ $1.8 \pm 0.5 \times 10^{-9}$ $3.9 \pm 0.4 \times 10^{-8}$ $1.6\pm0.2\times10^{-9}$ Initial rapid phase 4294 ± 1119 4330 ± 2443 t_{1/2} (Days) 3658 ± 99 493 ± 221 925 ± 328 390 ± 50 337 ± 36 337 ± 36 13 ± 2 Material Dissolved (%) 35.93 ± 4.05 0.47 ± 0.15 0.48 ± 0.20 3.29 ± 0.47 2.16 ± 0.13 2.08 ± 0.69 1.18 ± 0.18 0.47 ± 0.13 3.29 ± 0.47 PVP01 8650aR315 S850 R600 S750 S950 R204 R93

 a S650 is the same as R600.